10/773,414 8/29/04 SWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) s mixt. whereby two phases are obtained; (d) seps. the two phases tained; and (e) recovering the compd. I. The compds. I can be converted irbesartan which is a known angiotensin II receptor antagonist

L3 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2004:569369 CAPLUS COLONERY NUMBER: 141:225515 Synthesis 0? 2-butyl-3-[[2'-(1-trityl-1H-tetrazol-5-y]]biphenyl-4-y]]nethyl]-1, 3-diazaspiro[4,4]-non-ene-4-

one Nisnevich, Gennady: Rukhman, Igor: Pertsikov, Boris: Kaftanov, Julia: Dolitzky, Ben-zion Reparaceutical Industries Ltd., Israel: Teva Pharmaceuticals Uga, Inc. INVENTOR(S):

PATENT ASSIGNEE (S):

SOURCE: PCT Int. Appl., 27 pp. CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. DATE λl WO 2004072064
W: AE, AE, AG,
BG, BR, BR,
CU, CU, CZ,
ES, FI, FI,
15, JP, JP,
LK, LR, LS,
MZ, MZ, NA,
RW: BW, GB, GM,
GG, GH, CY,
MC, NL, FT,
GQ, GW, ML,
US 200424294
PRIORITY APPLN. INPO.: WO 2004072064 20040826 20040205 WO 2004-US3604

OTHER SOURCE(S):

CASREACT 141: 2255 \$5 2003-4659

Provided are 5 methods of making 2-butyl-3-[[2'-(1-trityl-1H-tetrazol-5-yl)bjhemyl-4-yl]methyl]-1,3-diazaspiro[4,4]non-1-ene-4-one [I], e.g. comprising the steps of: (a) reacting 1-(N'-pentanoylmanino )cyclopentanecarboxylic acid amide with 5-[4'-bromomethylbjhemyl -2-yl]-1-trityl-1H-tetrazole in the presence of an inorg, base, a solvent and a phase transfer catalystr (b) cooling the mixture; (c) adding water to

Search all reachons combining

1-(N'- pentanog () cyclopentane carboxylie acid amble
by RN (.e. all chemical
names for Hurrye)
and corresponding ester)

5-(4'-bromomothylbipliengl-2-yl)-1+rifil-14tehn ill.

1 het - this app.

Preparation of benzimidazole derivatives for

inhibiting neoplastic cells Sperl, Gerhard: Pamukcu, Rifat: Imkes, Ulrich: Piazza,

INVENTOR(S):

Gary A. Cell Pathways, Inc., USA PATENT ASSIGNEE(S): SOURCE:

U.S., 54 pp. CODEN: USXXXAM

DOCUMENT TYPE: LANGUAGE: Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 6348032 US 2002082280 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI US 1998-199094 US 2001-12672 US 1998-199094 B1 A1 20020219 19981123 20020627 MARPAT 136:183821

$$(R^3)_n \xrightarrow{N} R^2 \qquad 0 \xrightarrow{N} Me$$

$$(CR_2)_m - R^1 \quad I \qquad Br \qquad I$$

Title compds. I [wherein R1 = H, alkyl, benzenesulfonyl, or (un)substituted (hetero)aryl; R2 = H, (halo)alkyl, alkoxy(alkyl), (alkyl)amino, or carboxyl; R3 = halocarbonyl, carboxyl, haloalkylcarbonyl, alkoxycarbonyl, carboxyl, haloalkylcarbonyl, alkoxycarbonyl, aminosulfonyl, CN, (un)substituted carbamoyl, carbamoylalkyl, carbamoylalkenyl, or aryloxycarbonyl; n = 0-2; n = 0-2] where prepared for inhibiting neoplasia, particularly cancerous and precancerous lesions, without substantially inhibiting PGE-2. For example, a DMF solution of Et 3-butyrylamino-4-nitrobenzoate was added to NaOH (60% oil suspension) in a nitrogen environment at room temperature Dropwise addition of 2-bromobenzyl bromide ra

10 min span, followed by stirring for 1 h at room temperature and quenching

ice water, gave Et 3-{N-{2-bromobenzyl}butyrylamino}-4-nitrobenzoate.
Treatment with reduced Fe in AcOH and EtOH afforded the benzimidazole II.
Besides their utility as antitumor agents, I are also useful in the
treatment of diseases associated with abnormalities of cellular growth patterns such as benign prostatic hyperplasia, neurodegenerative diseases such as Parkinson's disease, autoimmune diseases including multiple solerosis and rheumatoid arthritis, infectious diseases uch as AIDS, and other diseases (no data).

REFERENCE COUNT: 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS

THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1993:22155 CAPLUS DOCUMENT NUMBER: 118:22155 TITLE: Preparation of substituted 1

118:22155
Preparation of substituted 1(2H)-isoquinolinones as angiotensin II antagonists
Patchett, Arthur A.; De Laszlo, Stephen E.; Greenlee, William J.
Herck and Co., Inc., USA
EUR. Pat. Appl., 68 pp.
CODEN: EFEXUM
Patent
English
1 INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. DATE DATE A1 19920909 GB, IT, LI, NL AA 19920907 A2 19930615 B4 19950419 EP 502575 R: CH, DE, FR, CA 2062211 JP 05148238 JP 07035372 PRIORITY APPLN. INFO.: EP 1992-200563 19920227 CA 1992-2062211 JP 1992-98999 19920303 19920306 US 1991-665491 US 1992-830621 A 19910306 A 19920211 OTHER SOURCE(S): MARPAT 118:22155

Title compds. I (R1 = (modified) H02C, H03S, NH202S, tetrazolyl, etc.; R2a, R2b = H, halo (di)(alkyl)amino, F3C, (substituted) aminosulfonyl, C1-6 alkyl, C1-6 alkowy, etc.; R3a = H, halo, C1-6 accylony, C3-7 cycloalkyl, etc.; R7a, F7b, R8a, R8b = H (substituted) C1-6 alkyl, piperarinyl, morpholino, etc.; R6 = (substituted) aryl, C1-6 alkyl, C2-5 alkynyl, etc.; E = bond, ininoalkylaulfonylalkylene, C0, etc.; R21 = H, halo, aryl, heteroaryl, etc.; r = substituted iterazolylcarbamoyl; x > bond, C0, O, s, etc.; are useful as angiotensin II antegonists (no data). Homophthalic anhydride in pyridine was added to valeryl chloride followed by treatment with NH4OH, and the mixture refluxed for 2 h to give 3-n-butyl-1(2H)-isoquinolinone. This was added to NHA and reacted with N-(triphenylmethyl)-5-[2-(4'-brommethylbiphenyl ]) tetrazol in DMF to give a product which was deprotected to give the title 3-butyl-2-[(2'-tetrazol-5-ylbiphen-4-yl)methyl]-1(2H)-isoquinoline. Pharmaceutical formulations comprising I are given. Claimed also are pharmaceuticals comprising I and antihypertensives, diuretics, angiotensin converting enzyme or Ca channel blocker.

L3 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

=> d 1 all

```
ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN 2004:696369 CAPLUS 141:225515
                             141:225515
Entered STN: 26 Aug 2004
Synthesis of 2-butyl-3-[[2'-(l-trityl-1H-tetrazol-5-yl)biphenyl-4-
yl]methyl]-1,3-diazaspito[4,4]-non-ene-4-one
Nisnevich, Gennady: Rukhman, Igor: Pertsitov, Boris: Kaftanov, Julia:
Bolitzky, Ben-zion
Teva Pharmaceutical Industries Ltd., Israel: Teva Pharmaceuticals Usa,
  TM
  PA
                         Inc.
PCT Int. Appl., 27 pp.
CODEN: PIXXD2
Patent
English
ICM CO7D403-10
28-10 (Meterocyclic Compounds (More Than One Hetero Atom))
Section cross-reference(s): 1
  50
                                                                                                                                                                                                                                 APPLICATION NO.
                             PATENT NO.
                                                                                                                                                                                                                                                                                                                                                       DATE
                                                                                                                            KIND DATE
PATENT NO.
                                                                                       CLASS PATENT FAMILY CLASSIFICATION CODES
       WO 2004072064 ICM C07D403-10 C07D403/10+257+235 US 2004242894 NCL 548/252.000
                             CASREACT 141:225515
                      ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
RL: CAT (Catalyst use): USES (Uses)
(catalysts for cyclocondensation of (peatancylamino
) cyclopentanecarboxanide with (bromeasthylbiphamyls)
) tetrazole; methods for prepn. of 2-butyl-3-[[2'-[1-trityl-]H-tetrazol-5-yl)) biphemyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)
64-18-6, Formic acid, uses 64-19-7, Acatic acid, uses 7647-01-0,
Hydrochoric acid, uses 1003-510-6, Hydrobromic acid, uses
RL: CAT (Catalyst use): USES (Uses)
(catalysts for imidation of valerimidate ester with
(aminomethylbiphemyly]) tetrazole; methods for preparation of
2-butyl-3-[[2'-(1-trityl-]H-H-tetrazol-5-yl) biphenyl-4-yl]methyl]-1,3-
diazaspiro[4,4]-non-ene-4-one)
57246-71-6, Westhyl valerimidate
RL: RCT (Reactant); RACT (Reactant or reagent)
(catalysts for imidation of valerimidate ester with
(aminomethylbiphemylyl) tetrazole; methods for preparation of
2-butyl-3-[[2'-(1-trityl-]H-H-tetrazol-5-yl) biphenyl-4-yl]methyl]-1,3-
diazaspiro[4,4]-non-ene-4-one)
76-05-1, Trifluorocaectic acid, uses
RL: CAT (Catalyst use): USES (Uses)
(catalysts for imidation of valerimidate ester with
5-(4'-aminomethylbiphenyl-2-yl)tetrazole; methods for preparation of
2-butyl-3-[[2'-(1-trityl-]H-tetrazol-5-yl) biphenyl-4-yl]methyl]-1,3-
diazaspiro[4,4]-non-ene-4-one)
999-09-7, Ethyl valerimidate 745814-12-4, Propyl valerimidate
745814-13-5, Butyl valerimidate 745814-12-4, Propyl valerimidate
745814-13-5, Pentyl valerimidate
RL: RCT (Reactant); RACT (Reactant or reagent)
(imidation of valerimidate ester with (aminomethylbiphenyl) tetrazole;
methods for preparation of 2-butyl-3-[[2'-(1-trityl-]H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)
638-29-9, Valeroyl chloride 1664-35-3, 1-Aminocyclopentanecarboxylic
acid ethyl ester 17193-28-1, 1-Aminocyclopentanecarboxylic
(methods for preparation of 2-butyl-3-[[2'-(1-trityl-]H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)
17450-51-2P, 5-(4'-Bromomethylbiphenyl-2-yl)-1-trityl-H-tetrazol-5-yl)biphenyl-4-y
                               /ASSI4-U9-99
RL: SPN (Synthetic preparation); PREF (Preparation)
  (methods for preparation of 2-butyl-3-{[2'-(1-trityl-1H-tetrazol-5-y]]biphenyl-4-y]l methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)
138402-11-6F, Irbesartan
```

RL: SPN (Synthetic preparation): THU (Therapeutic use): BIOL (Biological

```
AB Provided are 5 methods of making 2-butyl-3-[{2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro(4,4]non-1-ene-4-one (I), e.g. comprising the steps of: (a) reacting 1-(N'-pentanoylamino ) cyclopentanecarboxylic acid anide with 5-(4'-bromomethylbiphemyl 2-yl)-1-trityl-1H-tetrazole in the presence of an inorg, base, a solvent and a phase transfer catalyst; (b) cooling the mixture; (c) adding water to the mixture whereby two phases are obtained; (d) separating the two phases obtained; and (e) recovering the compound I. The compds. I can be converted to irbesartan which is a known angiotensin II receptor antagonist (blocker).

55 butyltrityltetrazolylbiphenylylmethyldiazaspirolenone prepn intermediate irbesartan

16 Ethers, uses

17 Riv NUU (Other use, unclassified); USES (Uses)

18 Riv NUU (Other use, unclassified); USES (Uses)

19 Riv Aliphatic, solvent; methods for preparation of

2-butyl-3-[(2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1, 3-diazaspiro[4,4]-non-ene-4-one)

17 Phosphonium compounds

18 CAT (Catalyst use); USES (Uses)

19 (catalyst for cyclocondensation of (pentanoylamino)

19 (cyclopentanecarboxamide with (bromomethylbiphemylyl)

10 (cyclopentanecarboxamide with (bromomethylbiphemylyl)

11 Cyclocondensation reaction

12 (cyclocondensation of (pentanoylamino) cyclopentanecarboxamide with (bromomethylbiphemylyl): Herterazol: sethods for preparation of 2-butyl-3-[(2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)

12 Acetaly

13 Riv NUU (Other use, unclassified); USES (Uses)

14 Acetaly

15 Riv NUU (Other use, unclassified); USES (Uses)

16 (Gylmes, solvent; methods for preparation of 2-butyl-3-[(2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)

14 Riv Auses

15 Riv NUU (Other use, unclassified); USES (Uses)

16 (Gylmes, solvent; methods for preparation of 2-butyl-3-[(2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)

17 Acetaly

18 (cyclopednsation reaction cataly
```

L3 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN

L3 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) study), PREP (Preparation), USES (Uses) (sethods for preps. of 2-butyl-3-[[2'-(1-trityl-1H-tetrazol-5-y1)biphenyl-4-y1]sethyl-1,3-diazaspiro[4,4]-non-ene-4-one)

IT 68-12-2, N,N-Dimethylformamide, uses 71-43-2, Banzene, uses 95-47-6, o-Xylene, uses 108-38-3, Toluene, uses 109-99-9, Tetrahydrofuran, uses 100-58-3, Toluene, uses 109-99-9, Tetrahydrofuran, uses 110-54-3, Hexane, uses 110-71-4, 1,2-Dimethoxyethane 119-64-2, Tetrahin 127-19-5, N,N-Dimethylacetamide 462-95-3, Diethoxymethane 164-04-4, Methyl tetr-butyl ether RL: NUU (Other use, unclassified), USES (Uses) (solvent, methods for preparation of 2-butyl-3-[2'-(1-trityl-1H-tetrazol-5-y1)biphenyl-4-y1]methyl]-1,3-diazaspiro[4,4]-non-ene-4-one)

### => FIL REGISTRY

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	22.35	22.56
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
CA SUBSCRIBER PRICE	ENTRY -2.92	SESSION -2.92

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\*

L4 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
RN 124750-51-2 REGISTRY
CN 1H-Tetrazole, 5-[4\*-(bromomethyl)[1,1\*-biphenyl]-2-yl]-1-(triphenylmethyl)(9C1) (CA INDEX NAME)
OTHER NAMES:
CN 4\*-Bromomethyl-2-(1-triphenylmethyltetrazol-5-yl)biphenyl
CN 4\*-Bromomethyl-2-(1-triyl-1H-tetrazol-5-yl)biphenyl
CN 4\*-Bromomethyl-2-(N-trityl-1H-tetrazol-5-yl)biphenyl
CN 4\*-Bromomethyl-2-(N-triphenylmethyl-1H-tetrazol-5-yl)biphenyl
CN 4\*-Bromomethyl-2-(N-triphenylmethyl-1H-tetrazol-5-yl)biphenyl
CN 4\*-Bromomethyl-2-(N-triphenylmethyl-1H-tetrazol-5-yl)biphenyl
CN 4\*-(8-Gromomethyl)biphenyl-2-yl)-1-trityl-1H-tetrazole
CN 5-[4\*-(Bromomethyl)biphenyl-2-yl]-1-(triphenylmethyl)-1H-tetrazole
CN 5-[4\*-(Bromomethyl)[1,1\*-biphenyl]-2-yl]-1-(triphenylmethyl)-1H-tetrazole
CN N-Triphenylmethyl-5-[2-[4\*-(bromomethyl)biphenyl)])tetrazole
CN N-Triphenylmethyl-5-[4\*-(bromomethyl)biphenyl-2-yl]tetrazole
CN [2\*-(1-Trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl bromide
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl bromide
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl)methyl bromide
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl)
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl)
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl)
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl)
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl]
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl]
CN [2\*-(Triphenylmethyl-1H-tetrazol-5-yl)biphenyl-4-yl]
C



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

194 REFERENCES IN FILE CA (1907 TO DATE)
194 REFERENCES IN FILE CAPLUS (1907 TO DATE)

#### => SET NOTICE LOGIN DISPLAY

NOTICE SET TO OFF FOR DISPLAY COMMAND SET COMMAND COMPLETED

=>

=> FIL REGISTRY

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	3.56	26.12
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	0.00	-2.92

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Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> S 177219-40-8/RN

L5 1 177219-40-8/RN

=> SET NOTICE 1 DISPLAY

NOTICE SET TO 1 U.S. DOLLAR FOR DISPLAY COMMAND

# SET COMMAND COMPLETED

=> D L5 SQIDE 1-

YOU HAVE REQUESTED DATA FROM 1 ANSWERS - CONTINUE? Y/(N):Y THE ESTIMATED COST FOR THIS REQUEST IS 6.15 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:Y

```
L5 ANSVER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
RN 177219-40-8 REGISTRY
CV Cyclopentanecarboxamide, 1-[(1-oxopentyl)amino)- (9CI) (CA INDEX NAME)
OTHER NAMES:
CN 1-(Pentanoylamino)cyclopentanecarboxamide
FS 3D COMCORD
MF C11 H20 N2 O2
CI COM
SR CA
LC STN Files: CA, CAPLUS, CASREACT, USPATFULL
UT.CA Caplus document type: Patent
RL.P Roles from patents: FREP (Preparation); RACT (Reactant or reagent)
```

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

4 REFERENCES IN FILE CA (1907 TO DATE) 4 REFERENCES IN FILE CAPLUS (1907 TO DATE)

#### => SET NOTICE LOGIN DISPLAY

NOTICE SET TO OFF FOR DISPLAY COMMAND SET COMMAND COMPLETED

=>

=> FIL REGISTRY

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	2.27	28.39
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	0.00	-2.92

FILE 'REGISTRY' ENTERED AT 10:01:33 ON 29 AUG 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 28 AUG 2005 HIGHEST RN 861926-07-0 DICTIONARY FILE UPDATES: 28 AUG 2005 HIGHEST RN 861926-07-0

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TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

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Structure search iteration limits have been increased. See HELP SLIMITS for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> S 745814-11-3/RN

L6 1 745814-11-3/RN

=> SET NOTICE 1 DISPLAY

NOTICE SET TO 1 U.S. DOLLAR FOR DISPLAY COMMAND

## SET COMMAND COMPLETED

=> D L6 SQIDE 1-

YOU HAVE REQUESTED DATA FROM 1 ANSWERS - CONTINUE? Y/(N):Y THE ESTIMATED COST FOR THIS REQUEST IS 6.15 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:y

L6 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
RN 745814-11-3 REGISTRY
CN Cyclopentanecarboxylic acid, 1-{(1-oxopentyl)amino}-, ethyl ester (9CI)
(CA INDEX NAME)
OTHER NAMES:
CN 1-(Pentanoylamino)cyclopentanecarboxylic acid ethyl ester
FS 3D CONCORD
NF C13 H23 N 03
SR CA
LC STN Files: CA, CAPLUS, CASREACT, USPATFULL
OT.CA Caplus document type: Patent
RL.P Roles from patents: RACT (Reactant or reagent)



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT \*\*

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

### => SET NOTICE LOGIN DISPLAY

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=>

=> d his

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FILE 'CAPLUS' ENTERED AT 09:48:37 ON 29 AUG 2005

L1 164 S BROMOMETHYLBIPHENYL?

L2 1044 S PENTANOYL?

L3 3 S L1 AND L2

FILE 'REGISTRY' ENTERED AT 09:58:37 ON 29 AUG 2005

L4 1 S 124750-51-2/RN

SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'REGISTRY' ENTERED AT 10:01:01 ON 29 AUG 2005

L5 1 S 177219-40-8/RN

SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'REGISTRY' ENTERED AT 10:01:33 ON 29 AUG 2005

L6 1 S 745814-11-3/RN

SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

=> fil caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
2.27 30.66

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL
TOTAL

CA SUBSCRIBER PRICE ENTRY SESSION 0.00 -2.92

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=> s L4 and (L5 or L6) 194 L4 4 L5 1 L6 L7 1 L4 AND (L5 OR L6)

=> d ibib abs

L7 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 2004:694369 CAPLUS
101:225515
Synthesis of 2-butyl-3=fft\*41\_trityl-1H-tetrazol-5yll biphenyl-4-yl] methyl]-1,3-diaza3piro[4,4]-non-ene-4one
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Teva Pharmaceutical Industries Ltd., Israel: Teva
Pharmaceuticals Usa, Inc.
PCT Int. Appl., 27 pp.
CODEN: PIOXID
COMMENT TYPE: Patent
LANGUAGE: Patent
LANGUAGE: English
PATENT INFORMATION:

			NO.			KIN	D	DATE APPLICATION NO.						DATE					
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WD 2004072064			A1 20040826			WO 2004-US3604						20040205							
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			15,	JP,	JP.	KE,	KE,	KG.	KG.	KP.	KP.	KP.	KR,	KR,	KZ,	ΚZ,	ΚŻ,	LC,	
			LK.	LR.	LS.	LS.	LT.	w,	LV.	MA,	MD.	MD.	MG.	MK.	MN.	KV,	MX,	MX,	
			MZ.	MZ.	NA.	NI													
		RW	BW.	GH,	GH,	ΚĖ,	LS.	MV.	MZ.	SD,	SL,	SZ.	TZ.	UG.	24.	ZV.	AT,	BE,	
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OTHER SOURCE(S):

Provided are 5 methods of making 2-butyl-3-[[2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-1,3-diazaspico[4,4]non-1-ene-4-one [1], e.g. comprising the steps of: (a) reacting 1-[N'-pentanoylamino]cyclopentanecar/borylic acid amide with 5-[4'-bromomethylbiphenyl-2-yl)-1-trityl-1H-tetrazole in the presence of an inorg, base, a solvent and a phase transfer catalyst; (b) cooling the mixture; (c) adding water to the mixture AB

L7 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) whereby two phases are obtained; (d) sepg. the two phases obtained; and (e) recovering the compd. I. The compds. I can be converted to irbesartan which is a known angiotensin II receptor antagonist (blocker).

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COST IN U.S. DOLLARS	SINCE FILE	TOTAL
·	ENTRY	SESSION
FULL ESTIMATED COST	3.10	33.76
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-0.73	-3.65

STN INTERNATIONAL LOGOFF AT 10:02:26 ON 29 AUG 2005